NIH -- W1 NA81

PAMELA GEHRON ROBEY

CSDB/NIDR/NIH Bldng 30 Rm 228 30 CONVENT DRIVE MSC 4320 BETHESDA, MD 20892

ATTN: SUBMITTED: 2001-11-19 16:19:04 PHONE: 301-496-4563 PRINTED: 2001-11-20 16:11:51

PHONE: 301-496-4563 PRINTED: 2001-11-20 TFAX: 301-402-0824 REQUEST NO.: NIH-10071782 E-MAIL: SENT VIA: LOAN DOC

5052780

NIH Fiche to Paper Journal

TITLE: NATURE

PUBLISHER/PLACE: Nature Publishing Group London:
VOLUME/ISSUE/PAGES: 1996 Jan 25;379(6563):311-9 311-9

DATE: 1996

AUTHOR OF ARTICLE: Lambright DG; Sondek J; Bohm A; Skiba NP; Hamm HE; Sigler PB

TITLE OF ARTICLE: The 2.0 A crystal structure of a heterotrimeric G

ISSN: 0028-0836

OTHER NOS/LETTERS: Library reports holding volume or year

0410462 8552184

SOURCE: PubMed
CALL NUMBER: W1 NA81
REQUESTER INFO: AB424

DELIVERY: E-mail: probey@DIR.NIDCR.NIH.GOV

REPLY: Mail:

NOTICE: THIS MATERIAL MAY BE PROTECTED BY COPYRIGHT LAW (TITLE 17, U.S. CODE)

-----National-Institutes-of-Health,-Bethesda,-MD------

The 2.0 Å crystal structure of a heterotrimeric G protein

David G. Lambright*†, John Sondek*, Andrew Bohm*, Nikolai P. Skiba‡, Heidi E. Hamm‡ & Paul B. Sigler*§

* The Department of Molecular Biophysics and Biochemistry, and the Howard Hughes Medical Institute, Yale University, 295 Congress Avenue, BCMM 154, New Haven, Connecticut 06510, USA

‡ Department of Physiology and Biophysics, University of Illinois at Chicago, Chicago, Illinois 60680, USA

The structure of a heterotrimeric G protein reveals the mechanism of the nucleotide-dependent engagement of the α and $\beta\gamma$ subunits that regulates their interaction with receptor and effector molecules. The interaction involves two distinct interfaces and dramatically alters the conformation of the α but not of the $\beta\gamma$ subunits. The location of the known sites for post-translational modification and receptor coupling suggest a plausible orientation with respect to the membrane surface and an activated heptahelical receptor.

Many extracellular signals are mediated by heptahelical receptors coupled to heterotrimeric guanine-nucleotide-binding proteins (G protein; subunits, $\alpha\beta\gamma$)¹⁻³. In response to diverse stimuli, including light, hormones and neurotransmitters, heptahelical receptors activate heterotrimeric G proteins by catalysing the exchange of GTP for GDP bound to the G, subunit. An agoniststimulated receptor activates as many as several hundred G proteins, which in turn activate a variety of downstream effectors including enzymes, ion channels4, and intracellular signalling pathways such as the MAP kinase cascade^{5,6}. In the resting state, the GDP form of the G_{α} subunit (G_{α} GDP) forms a high-affinity (nM) complex with the G_{β_0} heterodimer. Binding of the heterotrimeric complex to an agonist-activated receptor results in the release of GDP. Subsequent GTP binding disrupts the complex with the receptor and leads to the dissociation of $G_{\mu}GTP$ from $G_{\mu\nu}$ Both activated G_aGTP and the released G_{fr} subunits are free to interact with downstream components of the signalling cascade. As a consequence of an intrinsic GTP as activity in the G_x subunit, GTP is hydrolysed to GDP, thereby returning the system to its heterotrimeric resting state.

One of the best characterized heterotrimeric G-protein-coupled pathways is the visual cascade of the rod cell^{7,8}, where the light-activated heptahelical receptor, rhodopsin, couples to the heterotrimeric G protein transducin $(G_{ta\beta\gamma})$. G_{tz} GTP activates a potent cyclic GMP phosphodiesterase by displacing its inhibitory γ -subunits. A comparison of the structure of G_{tz} complexed with either GDP⁹ or GTP- γ S¹⁰ (a non-hydrolysable GTP analogue) revealed the nucleotide-dependent structural changes that distinguish the GTP-bound (active) form from the GDP-bound (inactive) form. The structural changes occurring within the G_z subunit are not global, but rather are restricted to three adjacent 'switch' regions. Highly localized rearrangements within the context of an otherwise active GTP-bound conformation stabilize the transition state for GTP hydrolysis^{11,12}.

We now describe the refined 2.0 Å crystal structure of a heterotrimeric G-protein complex. A comparison with the previously determined $G_{ta}GDP$ structure⁹ and that of free $G_{t\beta}$, (ref. 26) establishes the conformational changes that occur in each subunit to form the heterotrimer interface and hence reveals the mechanism of GTP-induced release and activation of G_{β} . The sites of post-translational modification on G_{tx} and G_{ty} combined

with the receptor-binding regions of G_{ts} , deduced from biochemical studies, occur on a common face that orients the heterotrimer with respect to the membrane surface.

Structure determination and refinement

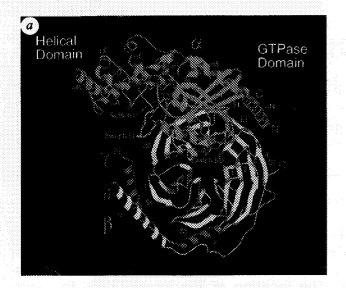
A modified heterotrimeric complex of transducin was prepared by mixing endo-Lys-C-proteolysed $G_{t\mu}$, with a G_{tx}/G_{tz1} chimaera $(G_{tx/tz1})$ in which residues 216–294 of G_{tx} were replaced with the corresponding residues (220–298) from G_{tz1} (ref. 13). This chimaera binds guanine nucleotides, assumes an active conformation in the presence of aluminium fluoride, and interacts with $G_{t\beta\gamma}$ and rhodopsin in a light-dependent manner¹³. Proteolysis of $G_{t\beta\gamma}$ with endo-Lys-C removes only the farnesyl modification and three residues from the C terminus of $G_{t\gamma}$ while leaving $G_{t\beta}$ intact. $G_{tx/tz1}$ was expressed in *Escherichia coli* as a soluble fusion with an N-terminal 6xHis tag which was not removed before crystallization. Diffraction-quality crystals of the heterotrimeric complex in the monoclinic space group C2 were grown as described (Table 1).

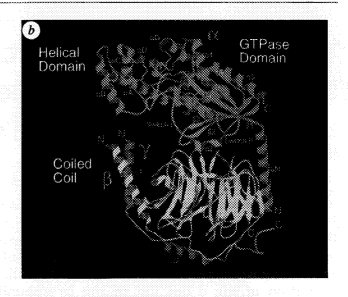
The structure was solved at 3.0 Å by multiple isomorphous replacement (MIR) using isomorphous differences from a Hg(OAc), derivative and isomorphous as well as singlewavelength anomalous differences from a CH3HgCl derivative and a selenomethionine (SeMet) derivative substituted in the G_{ta/at} subunit. The intial map was improved with solvent flattening and histogram matching. A molecular replacement (MR) solution for the G_{ta/in1} subunit was obtained at 4Å using the G_{ta}GDP structure as a search model. Although the phases obtained from the MR solution were not sufficiently accurate to generate an interpretable map of the protein, they were useful for calculating derivative difference Fourier maps which allowed the location of heavy atom sites in each of the derivatives. Using Sigma A weights to reduce model bias, an interpretable map was calculated at 3.0 Å resolution with phases derived from multiple isomorphous replacement and anomalous scattering combined with phases from the partially refined molecular replacement model.

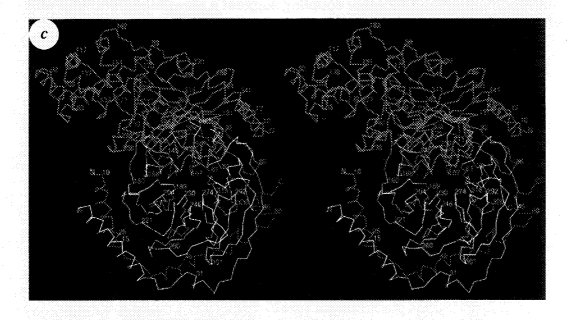
The model was refined with simulated annealing against the SeMet intensities from 6.0 to 2.0 Å resolution (Table 1). The present model includes residues 6–343 for G_{t_2,t_3} , residues 2–340 for G_{t_2,t_3} , residues 8–66 for G_{t_2,t_3} a GDP molecule, 630 water molecules, and has an R value of 20.7% for all data (19.6% for data $>2\sigma$) and a free R value of 29.5% (28.4% for data $>2\sigma$) for a randomly selected subset (10%) of the data omitted before the start of refinement. An example of the $2F_o - F_c$ electron density map is shown in Fig. 2d. The structure has

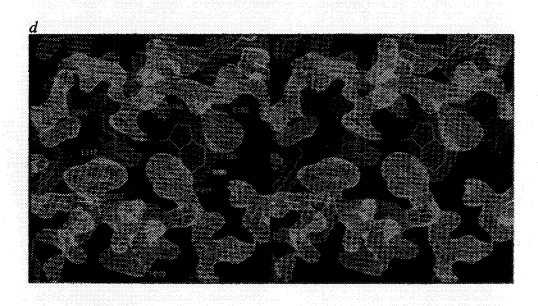
[†] Present address: Program in Molecular Modicine, University of Massachusetts Medical Center, 373 Plantation Street, Worcester, Massachusetts 0,605, USA. \$To whom correspondence should be addressed.

ARTICLES









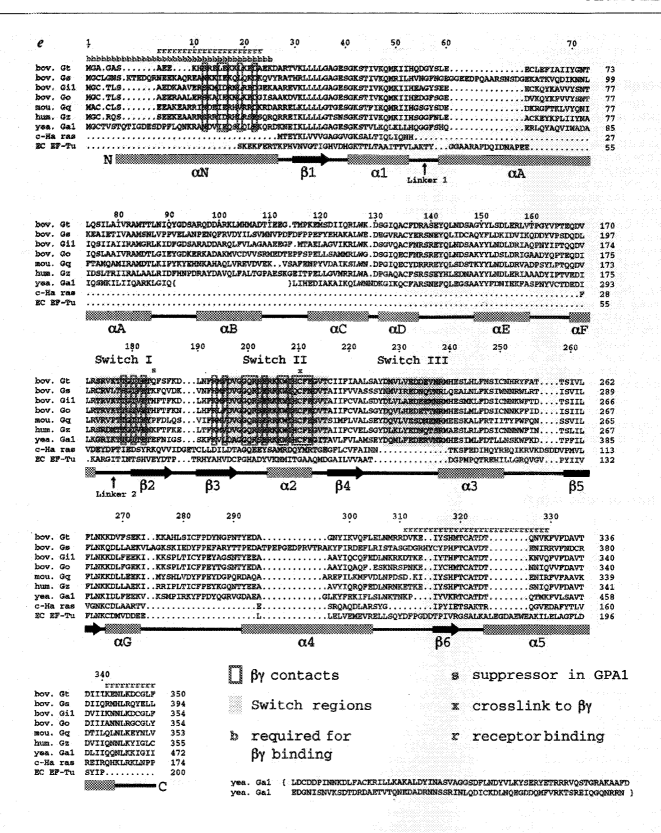


FIG. 1 Structure of the $G_{\text{to/nz}}G_{\text{to/n}}$ heterotrimeric complex with $G_{\text{to/nz}}$ shown in green, $G_{\text{to/n}}$ in yellow, $G_{\text{to/nz}}$ in red, GDP in magenta, and the switch I–III regions of $G_{\text{to/nz}}$ in cyan. a, Ribbon drawing of the complex viewed down the axis of the β -propeller domain of $G_{\text{to/nz}}$ b, Ribbon drawing of the complex rotated 70° about the horizontal axis compared with the view in a. c, Stereo pair showing the $C\alpha$ atoms of the complex oriented as in a. d, $2F_{\text{obs}}-F_{\text{calc}}$ electrondensity map contoured at 1.5σ showing residues in the switch interface.

e, Sequence alignment for G_π subunits with the secondary structure corresponding to G_{tolar} shown below the aligned sequences. Residues that contact G_{tolar} are indicated with red boxes and residues in the switch regions are highlighted in light blue. Also indicated are residues that have been implicated as in or near the G_{β_0} and receptor binding regions based on biochemical experiments and yeast genetics.

excellent stereochemistry and ϕ - ψ angles within allowed regions of the Ramachandran plot. A more detailed description of the structure solution and refinement is given in the legend of Table 1.

Overall structure

The structure of the heterotrimeric $G_{t\alpha/i\alpha}GDP\cdot G_{t\beta\gamma}$ complex is depicted in Fig. 1. The $G_{t\alpha}$ subunit has three distinct structural components: (1) a Ras-like GTPase domain consisting of a sixstranded β -sheet surrounded by six helices ($\alpha 1 - \alpha 5$ and αG); (2) an entirely α -helical domain consisting of a long central helix (α A) surrounded by 5 shorter helices ($\alpha B - \alpha F$); and (3) an N-terminal helix (αN) that projects away from the remainder of the $G_{t\alpha}$ subunit. The guanine nucleotide binds tightly in a deep cleft between the GTPase and helical domains. In general, the helical and GTPase domains as well as the guanine nucleotide closely resemble their counterparts in G_{tt}GDP, except for significant changes where the GTPase domain contacts G_{tg}. The N-terminal helix could not be observed in previously determined G_{tx} structures⁹⁻¹¹ where it was removed by proteolysis and was disordered in the structure of $G_{iz1}GTP\gamma S^{12}$. The conformation of the Nterminal helix does not resemble that observed in the crystal structure of G₁₂₁GDP¹⁴.

The $G_{t\beta}$ subunit has an N-terminal helix followed by a repeating module of seven similar β -sheets, each with four antiparallel strands, that form the 'blades' of a β -propeller structure (Fig. 1a). The outside strand of each sheet, together with the inner three strands of the following sheet, comprise the structural unit corresponding to the 'WD'-sequence repeat first noted in $G_{t\beta}^{15}$. The $G_{t\gamma}$ subunit contains two helices and has no inherent tertiary structure. The N-terminal helix of $G_{t\beta}$ forms a coiled-coil with the N-terminal helix of $G_{t\beta}$ whereas the remainder of $G_{t\gamma}$ interacts extensively with the β -propeller domain of $G_{t\beta}$. The details of the $G_{t\beta\gamma}$ structure are described in an accompanying paper²⁶.

\mathbf{G}_{α} interactions with $\mathbf{G}_{\beta\gamma}$

As shown in Fig. 2a, the interaction between the G_{tx} and $G_{t\beta\gamma}$ subunits occurs at two distinct interfaces. The most extensive interface involves residues in, or adjacent to, the switch I and II regions of G_{tx} that interact with residues from loops and turns at the top of the β -propeller domain of $G_{t\beta}$. The second interface is formed between the N-terminal helix of G_{tx} and the side of the 'switch interface' because it involves regions of G_{tx} that have been shown previously to undergo nucleotide dependent conformational changes and the latter as the 'N-terminal interface' because it involves only the N-terminal helix of G_{tx} .

The switch interface buries $\sim 1,800 \, \text{Å}^2$ of solvent-accessible surface area and includes residues from $\beta 2$, $\beta 3$, and $\alpha 2$ in $G_{t\alpha}$ as well as residues from five of the seven sheets in $G_{t\beta}$. Fig. 2b shows the core of this tightly packed interface which consists of the side chains from six hydrophobic residues in $G_{t\alpha}$ (Ile 180, Phe 195, Trp 207, His 209, Cys 210 and Phe 211) and four hydrophobic residues in $G_{t\beta}$ (Tyr 59, Trp 99, Met 101 and Leu 117). The interaction is further stabilized by a network of hydrophilic interactions including both main-chain and side-chain hydrogen bonds and is flanked by two ion pairs. The specific interactions are shown schematically in Fig. 2g.

In contrast, only $\sim 900\,\text{Å}^2$ of solvent-accessible surface area is buried in the N-terminal interface. As depicted in Fig. 2c, five residues on one side of the N-terminal helix of G_{tx} (Ser 12, Glu 16, Leu 19, Asp 22 and Ala 23) interact with four residues from the first sheet of G_{ty} (Leu 55, Lys 78, Ile 80, Lys 89). The conserved Lys 89 plays an essential role through a tripartite interaction in which its backbone NH donates a hydrogen bond to the hydroxyl group of Ser 12 while its methylene groups pack against the side chains of Leu 15 and Leu 19 thereby positioning its amino group to form an ion pair with the carboxylate group of Glu 16. In addition to enhancing the stability of the complex, the N-terminal interface also determines the location of the modified N terminus of G_z .

In contrast to extensive interactions with G_{tg} , no direct interactions are observed between G_{tx} and G_{ty} . However, the first five residues of G_{tx} as well as the first eight and last four residues of G_{ty} are disordered. Therefore, the absence of interactions between G_{tx} and G_{ty} in the present structure does not rule out the possibility that such interactions might occur under a more physiological setting. However, the extent of such an interaction would be limited.

Numerous biochemical and mutational studies have implicated the N terminus and switch II regions of G, in the interaction with G_{β_i} . Limited proteolysis 16, a monoclonal antibody directed against the N terminus^{17,18}, and expression of N-terminally truncated G_a subunits^{19,20} indicate that the myristoylated N terminus and some portion of the first 25 residues of G, are essential for high-affinity binding to $G_{\rho,r}$. The ability of $G_{o\alpha}$ to be ADP-ribosylated at Cys 351 by pertussis toxin is impaired by mutations at residues corresponding to Lys 17 and Lys 31 in G_{1x}^{21} . Both residues are exposed to solvent and neither contacts $G_{t\beta}$, suggesting that they may contribute to interactions with the toxin and/or the negatively charged membrane surface. The Gly 226 to alanine mutation in G_{sx} that disrupts the GTP-induced conformational changes in the switch II region also blocks the release of $G_{\beta_1}^{22,23}$, consistent with the extensive interaction between $G_{t\beta}$ and the switch regions of $G_{t\alpha}$. Finally, the conserved Cys 215 in the switch II region of G_{ox} can be crosslinked by a 15 A divalent alkylating agent to either of two conserved cysteines in $G_{t\beta}$ (Cys 204 and Cys 271)²⁴. The crosslinks are compatible with the location of these cysteine residues in the structure (see Fig. 2e).

Regions of G_{β} that contact G_{α} have been implicated by yeast genetics. Mutational changes in the yeast mating factor G_{β} (ste4p) at residues corresponding to Trp 99 and Met 101 in Gtx (Fig. 2e) cause constitutive activation of the pheromone signalling pathway and are defective for interaction with the yeast mating factor G_a (GPA1) in the yeast two-hybrid assay²⁵. Strikingly exposed on the surface of free $G_{t\beta y}$, these residues lie at the core of the hydrophobic switch interface with $G_{t\alpha}$ in the heterotrimeric complex (Fig. 2b). In addition, some of the constitutively active mutants in the yeast G_{θ} can be suppressed by a second site revertant (E307K)²⁵ at a residue corresponding to Gln 184 in G_{to}, which lies at the edge of the switch interface. Inspection of the heterotrimer structure predicts that a lysine residue at this position would be situated to form an ion pair with a non-conserved aspartic acid residue (Asp 133 in yeast $G_{\scriptscriptstyle B}$) located at the position of Arg 96 in G_{tB}.

Most of the contacts involve residues that are highly conserved in both G_{α} (Fig. 1e) and G_{β} (see accompanying paper²⁶). Substitutions at these positions are typically conservative even in sequences from distantly related organisms. It is therefore likely that the interactions observed here can be generalized to other members the heterotrimeric G-protein family. Moreover, the structure does not reveal an obvious basis for specificity in the pairing of particular G_{α} and $G_{\beta\gamma}$ subunits. Thus, the extent to which specificity is determined at the level of the G_{α} interaction with $G_{\beta\gamma}$ or at the level of the heterotrimer interaction with the receptor remains an interesting question.

Conformational changes in G

Figure 3 shows a superposition of the previously determined structures of free $G_{tx}GDP^9$ and $G_{tx}GTP\gamma S^{10}$ onto that of $G_{tx}GDP$ in the heterotrimeric complex. The overall conformation is preserved except for residues within the switch I and II regions and a few flexible loops. The differences between the free and heterotrimeric forms of $G_{tx}GDP$ are most pronounced for residues that interact with $G_{t\beta}$ directly. Some of these are critically conserved residues that are important to the structure and function of G_xGTP , such as Gly 199 which triggers conformational changes in the switch II region through an interaction with the γ phosphate in the $GTP^{9,10}$, and Gln 200 which stabilizes the transition state for GTP hydrolysis 11,12 .

The most dramatic changes occur in the switch II region. In free

 $G_{tz}GDP$, the $\beta 3$ strand terminates in a tight turn (Gly 198–Gln 200) that precedes the $\alpha 2$ helix (Arg 201 to His 209). As opposed to the taut and compact GTP-bound form, the $\alpha 2$ helix in free $G_{tz}GDP$ is a loosely configured 3_{10} helix with conserved hydrophobic and polar residues exposed and accessible to proteases. In order to engage the $G_{t\beta}$ subunit, the residues from Gly 198 to Ser 202 adopt an extended conformation, allowing the

remainder of the $\alpha 2$ helix to swing outwards, exposing and positioning residues to interact with $G_{1\beta}$. As a consequence, the hydrogen-bonding network between the $\beta 2$ and $\beta 3$ strands is extended at the expense of that between $\beta 3$ and $\beta 1$. These changes do not significantly alter the solvent-accessibility of the nucleotide. However, it is likely that $G_{\beta \gamma}$ binding stabilizes the otherwise flexible switch I and II regions, perhaps accounting for the

		TABLE :	Structure de	eterminatio	n and refinem	ent			
Crystal	Source			ta collectio ion limit (Å		R _{sym} (%)*		% Complete
Native SeMet MeHgCl Hg(OAc) ₂	R-axis II NSLS X25 R-axis II Xentronics	2.5 2.1 3.5 3.4				7,2 6.3 9.1 11.	1 1		97 99 85 87
			MIR p	hasing stat	stics				
Resolution limit (Å) Phasing power†‡ SeMet SeMet (anomalous) Hg(OAc) ₂ MeHgCl MeHgCl (anomalous)	0.32 0.21 1.14 1.42 0.05	10.17 0.63 0.24 1.71 1.58 0.27	7.27 0.54 0.30 1.90 1.79 0.36	5.66 0.53 0.32 2.33 2.41 0.30	4.63 0.38 0.28 1.67 1.93 0.23	3.92 0.28 0.28 1.16 1.34 0.15	3.40 0.23 0.25 0.95 0.95 0.07	3.00 0.23 0.21	Overall 0.30 0.26 1.42 1.55 0.16
Mean figure of merit	0.43	0.57	0.56	0.54	0.46	0.36	0.23	0.13	0.31
			Rot	tation searc	h		Dъ	fined PC (Ω)	. 2
		Euler angles				Highest			Highest
Search model	Θ_1	$oldsymbol{\Theta_2}$			Θ_3	peak		false peak	
G _{tx} GDP	29.3		88.7		42.7	42.7 15.7 σ			5.5σ
			Tran	slation sea	rch				
Canada manua di di		Fractional coordinates				Highest		ation function %§ Highest	
Space group C2	<i>x</i> 0.21		у 0.00		2 0.15		peak 21σ		false peak 16σ
02	0.21		0.00		0.13		210		100
	Resolution	on (Å)	Refinement R factor		Free R factor		ctor	No. of reflections	
Data with $F > 2\sigma$ All data R.m.s. deviations	6-2. 6-2.		19.6 20.7 Bond lengths 0.01			28.4 29.5 Å Bond angles 1.5°			49,438 54,174

Gize, was isolated from photolysed bovine retinal rod outer segments (ROS) by selective extraction with GTP49 and the subunits separated by chromatography on Cibracon blue Sepharose as described⁹. G_{ip} , was proteolysed with 10 U of the lysine-specific protease endo-Lys C for 12 hours at 4 °C followed by cation exchange chromatography on Mono-Q (10 mM Tris, pH 7.5, 0.1–0.2 M NaCl gradient over 30 column volumes). The $G_{tz/tz}$ chimaera was expressed as a soluble 6xHis fusion in *E. coli* as described¹³. Diffraction-quality crystals of the heterotrimeric complex were grown in microseeded hanging drops containing 10 mg ml⁻¹ protein, 10% PEG-8000, 50 mM Tris, pH 8.0, 10% glycerol, 50 mM NaCl and 0.1% β-mercaptoethanol over a well containing the identical solution without the protein. Crystals appeared in 1–2 days and grew to a maximum of 0.3 \times 0.3 \times 0.5 mm in 1 week. The space group is C2 with unit cell dimensions of a=133.4, b=91.4, c=83.2 Å. Crystals were incubated for 12 h at 4 °C in a cryoprotectant stabilizer solution containing 30% PEG-8000, 10% glycerol, 50 mM Tris, pH 8.0. The Hg2+ and CH3Hg+ derivatives were obtained by soaking in the presence of 0.1 mM mercuric acetate and 1 mM CH₃HgCl respectively. Crystals were flash-frozen in liquid propane which was cooled with liquid nitrogen. The native and CH₃Hg⁺ derivative data sets were collected at $-150\,^{\circ}$ C on an R axis II image plate system and processed with DENZO (Z. Otwinowski). The Hg²⁺ derivative data set was collected at 100K on a Xentronics area detector and processed with XDS⁵⁰. Diffraction data for the Se-methionine derivative were collected at 100K at the peak of the Se absorption edge at the X-25 beamline of the National Synchrotron Light Source at Brookhaven, processed with DENZO (Z. Otwinowski). All data sets were scaled with SCALEPACK (Z. Otwinowski). The location of the Gts/le1 subunit was found by molecular replacement using as a search model the Gts/GDP structure in which residues in the range of the chimera were truncated to alanine. A rotation search using Patterson correlation refinement⁵¹ resulted in a single solution. A translation search yielded a clear solution with an R value of 50.5% after rigid-body refinement. Heavy-atom sites were located from isomorphous and anomalous differences Fouriers with phases calculated from the molecular replacement solution for the G_{ta} subunit. The coordinates and occupancies of the heavy atoms were refined with ML-PHARE. Initial MIR phases were improved by solvent flattening, histogram matching, and partial model combination using DPHASE (G. van Duyne). All computations for molecular replacement and subsequent refinement were carried out with X-PLOR 3.1 (ref. 52). Interactive model building was done with O (ref. 53).

* $R_{\text{sym}} = \Sigma |I_h - \langle I_h \rangle |\Sigma I_h$

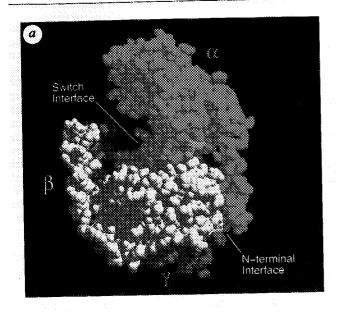
†Phasing power = $\Sigma |F_H|/\Sigma |F_{PH}obs| - |F_{PH}calc|$.

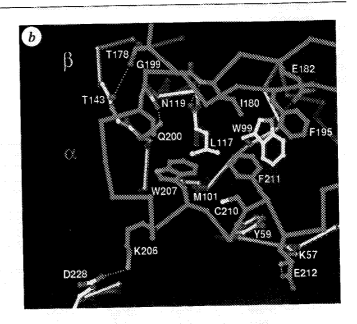
‡ Anomalous phasing power = $\Sigma |F''_{H}|/\Sigma |ADobs| - |ADcalc|$.

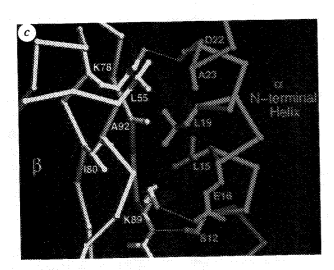
$$\label{eq:PCO} \S\,\%\text{PC}(\Omega) = \frac{\left\langle |E_{\text{obs}}|^2|E_{\text{m}}(\Omega)|^2 - \left\langle |E_{\text{obs}}|^2|\right\rangle \left\langle |E_{\text{m}}(\Omega)|^2\right\rangle \right\rangle}{\left[\left\langle |E_{\text{obs}}|^4 - \left\langle |E_{\text{obs}}|^2\right\rangle^2\right\rangle \left\langle |E_{\text{m}}(\Omega)|^4 - \left\langle |E_{\text{m}}(\Omega)|^2\right\rangle^2\right\rangle\right]^{1/2}},$$

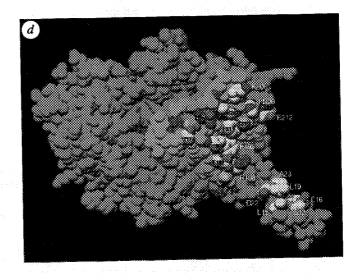
where E_{obs} denotes the normalized observed structure factors and E_m denotes the normalized structure factors for the search model place in a triclinic unit cell with geometry identical to that of the crystal³³.

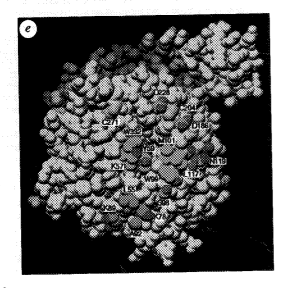
ARTICLES

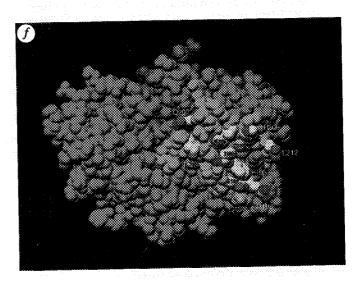












NATURE - VOL 379 - 25 JANUARY 1996

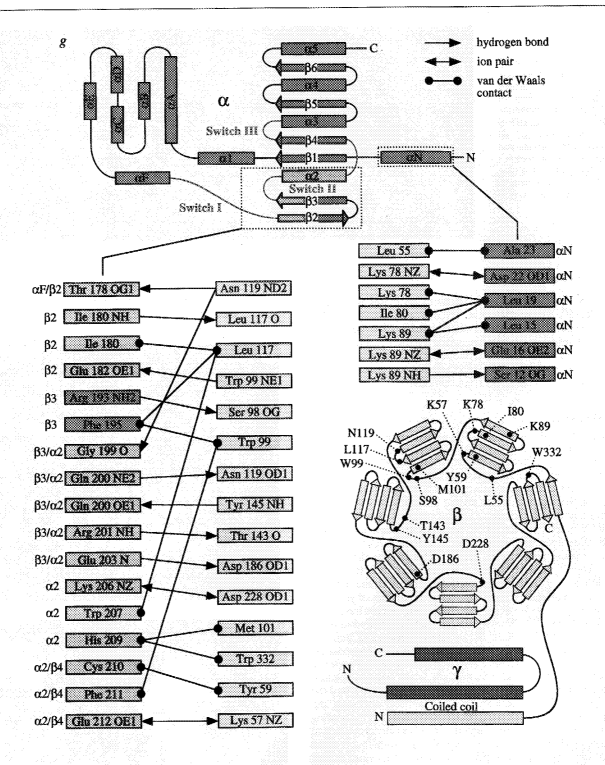
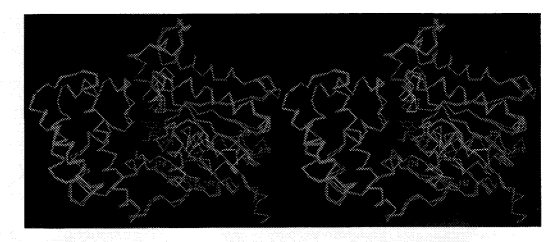


FIG. 2 Contacts in the two independent heterotrimer interfaces. a, Space-filling model showing the two distinct interfaces between the G_{tz} and $G_{t\bar{p}_1}$ subunits. b, Ball-and-stick model of the switch interface showing α -carbons (green G_{tz} ; yellow in $G_{t\bar{p}}$) and selected side chains (green in G_{tz} and yellow in $G_{t\bar{p}}$, except for oxygens and nitrogens which are red and blue, respectively). c, Ball-and-stick model of the N-terminal interface showing α -carbons and selected side chains coloured as in a. d, Space-filling model of $G_{tz/tz}GDP$ (green) showing the switch I and II regions (cyan) and the residues that contact $G_{t\bar{p}}$ (yellow, except oxygens and nitrogens which are red and blue, respectively). View is similar to the canonical orientation described previously $^{9-11}$. Residues identified in biochemical or mutational studies are highlighted in orange. e, Space-filling model of $G_{t\bar{p}_1}$ (yellow) showing the residues that contact $G_{tz/tz}GDP$ (green, except oxygens and nitrogens which are red and blue, respectively) in the switch interface. View is rotated by about the y-axis compared to d. Residues corresponding to Trp 99 and

Met 101 have been implicated by yeast genetics as essential for the interaction between $G_{\rm a}$ and $G_{\rm \beta r}$. Two cysteines in $G_{\rm \beta}$ (Cys 204 and Cys 271) that can be chemically crosslinked to Cys 215 in $G_{\rm ox}$ (Cys 210 in $G_{\rm to}$) are indicated in orange. Cys 210 lies at the edge of the switch II interface and makes van der Waals contact with Tyr 59 in $G_{\rm to}$. At a distance of 13 Å from Cys 210, Cys 204 is located in a turn between the second and third strands of the fourth WD repeat and make van der Waals contact with the methylene groups of Lys 206 in $G_{\rm to}$. Cys 271, on the other hand, occurs in the loop connecting the outer and inner strands of the sixth WD repeat and lies at a distance of 18 Å from Cys 210 and 10 Å from edge of the switch interface. $f_{\rm c}$ Space-filling model of $G_{\rm to}$ GTP-,S (coloured and oriented as in d) showing how the switch interface is disrupted by the GTP-induced conformational changes. $g_{\rm c}$ Diagram of the interactions between $G_{\rm to/to}$ and $G_{\rm to}$ in the heterotrimeric complex.

FIG. 3 Structural changes in Gt. and $G_{t \beta_i}$ subunits. Stereo pair showing the least-squares superimposed Ca traces of free $G_{t_{\alpha}}GDP$, heterotrimeric G, GDP, and $G_{tx}GTP_{\gamma}S$ oriented as in Fig. 1a. All three molecules are coloured green except for the switch I and II regions, which are blue for free GtaGDP, red for heterotrimeric $G_{tx/\alpha}GDP$, and yellow for GtaGTP7S.

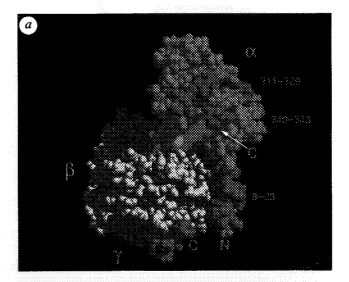


observation that the interaction with $G_{\beta\gamma}$ decreases the rate of GDP release²⁷. On replacement of GDP with GTP, local but dramatic conformational changes disrupt nearly all of the contacts in the switch interface, thereby triggering the disengagement of G_a and G_{θ_0} . It is also likely that the structure of the N-terminal helix of G_n depends on the interaction with G_n , as its conformation in the heterotrimer differs markedly from that observed in the GioDP structure14.

Implications

In contrast to the dramatic structural reorganization in the switch I and II regions of Gw significant structural changes are not observed in $G_{t\theta y}$. Although the structures of $G_{t\theta y}$ alone and bound to G_{tx} were independently determined, the r.m.s. deviation in C α positions after superposition is only 0.72 Å. Thus, the $G_{t\beta\gamma}$ subunits appear to function as a rigid unit with critical residues prepositioned to interact with GtaGDP or other signalling components. Whereas G_x activation proceeds through nucleotidedependent structural reorganization, activation of $G_{\beta\gamma}$ occurs solely as a consequence of its release. This implies that the G_{α} subunit acts as a negative regulator of G_{β_i} by restricting its degrees of freedom and/or masking sites on the surface of G_{β} that interact with downstream signalling components. The regions of yeast G_{β} , that interact with downstream signalling components have been mapped genetically^{54,57}. None of the residues implicated in these studies lie within either of the two interfaces between G_a and G_{ac} Instead, they map to the coiled-coil and a region immediately adjacent to the switch interface (Fig. 3 of accompanying paper²⁶). However, as both G_{α} and effectors are large macromolecules, steric constraints will extend the masked region of G₈ beyond the actual van der Waals contact surface, thereby extending the effective 'footprint' of G_{α} on $G_{\beta\gamma}$. Thus the G_{α} and effector binding sites on G_{R_2} need not overlap directly. In this respect, the helical domain of G_{α} (Fig. 1b) appears to be positioned such that it could sterically hinder the access of effectors that bind to regions similar to those implicated for yeast G_{β_i} .

Both the G_a and G_b, subunits undergo post-translational modification, resulting in the covalent attachment of specific lipophilic groups. The G_{γ} subunits are farnesylated $(G_{\gamma 1})^{28,29}$ or geranyl geranylated $(G_{\gamma 2-4})^{30-32}$ at the cysteine in the C-terminal CAAX motif. Several G_{α} subunits are subject to N-terminal myristoylation $(G_{12}, G_{02}, G_{22})^{33}$ or heterogeneous acylation $(G_{12})^{34,35}$ at Gly 2 and most (except G_{12}) are regulated by reversible palmitoylation at Cys $3^{36,37}$. These modifications are essential for membrane association. In addition, myristoylation of G_{12}^{38} and membrane association. In addition, myristoylation of G_{α}^{38} and prenylation of G, increase the stability of the heterotrimer complex³⁹. Experiments with G_{tx} indicate that this stabilization occurs only in the presence of membranes or detergent micelles, suggesting that the primary role of the lipophilic modifications is to co-localize and orient the subunits on the membrane surface⁴⁰.



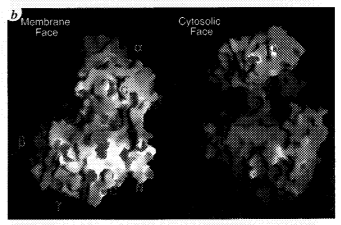


FIG. 4 Face of the heterotrimeric complex that is proposed to interact with the membrane surface and photoactivated rhodopsin. a, Space-filling model highlighting the sites of hydrophobic post-translational modification (N terminus of G, and C terminus of G,) and regions that are known from biochemical experiments to contact receptors (purple). The orientation is rotated about the vertical axis by $\sim\!70^\circ$ with respect to the view in Fig. 3 b. Solvent-accessible surface coloured according to electrostatic po using GRASP⁵⁶. The electrostatic potential is contoured in the ran $-10 k_B T$ (red) to $+10 k_B T$ (blue) where k_B is Boltzmann's constant T (blue) where k_s is Boltzmann's constant e (K). NATURE · VOL 379 · 25 JANUARY 1996 the absolute temperature (K).

Although the G_{α} and G_{γ} subunits in this study lack these modifications, their sites of attachment can be approximated from the location of the unmodified N- and C-termini of G, and G respectively. In the present structure, the first four residues (Gly-Ala-Gly-Ala) from the N terminus of G₁₂ and the last four residues (Lys-Gly-Gly-Cys) from the C terminus of G_t, are not observed. The amino-acid sequences suggest that both regions could be conformationally flexible in the absence of other interactions. Moreover, the structures of $G_{t\omega}$ and $G_{t\beta\gamma}$ do not reveal obvious binding sites for large hydrophobic groups. Significantly, the observed N terminus of $G_{t\alpha}$ and the observed C terminus of $G_{t\alpha}$ are located 18 A apart on a common face of the heterotrimer. Thus, the structure is consistent with the hydrophobic modifications inserting simultaneously and cooperatively into the lipid bilayer rather than binding to sites on the protein and also supports the conclusion that the enhanced stability of the heterotrimer complex results from co-localization of the subunits to membranes or detergent micelles.

The receptor binding surface of G_{α} has been mapped approximately by biochemical and mutational data. Many experiments indicate that the C terminus of G_{α} is an essential region for receptor coupling. ADP ribosylation by pertussis toxin⁴¹, mutations^{42,43} and peptide-specific antibodies⁴⁴, all directed at the C terminus, uncouple G proteins from receptors. Experiments with G_{α} chimaeras indicate that this region is also important in determining G-protein receptor specificity⁴⁵. A peptide corresponding to the C-terminal 11 amino acids has been shown to interact directly with photoactivated rhodopsin^{46,47}. In addition, the C-terminal peptide and peptides corresponding to the end of the $\alpha 4/\beta 6$ loop through the beginning of $\alpha 5$ (residues 311–329) as well as most of the N-terminal helix (residues 8-23) inhibit G_t binding to rhodopsin⁴⁷.

The approximate locations of the sites of post-translational modification and receptor coupling occur on a common face of the heterotrimeric complex (Fig. 4a). This side of $G_{t\beta\gamma}$ is surprisingly flat, and electrostatic calculations reflect a predominantly

neutral surface interspersed with patches of positive charge, in contrast to the other faces where the electrostatic potential is dominated by negatively charged residues (Fig. 4b). Thus, a plausible model would orient this face of the heterotrimer against the negatively charged membrane surface allowing the hydrophobic modifications to insert simultaneously into the lipid bilayer while positioning the known receptor contact sites of G_{α} to interact with an activated heptahelical receptor. In addition, $G_{\iota\beta_7}$ has also been shown to interact directly with rhodopsin⁴⁸; however, the regions that contact the receptor have not been identified. One possible receptor contact site on $G_{i\theta}$, consistent with the location of the approximate receptor contact regions of Ga (Fig. 4a), is the region adjacent to the switch and N-terminal interfaces. The observation that G_{θ_2} interacts with both the receptor and the switch I/switch II region of G_x , which flanks the nucleotide binding site and buttresses the critical phosphate binding loop, gives stereochemical support to a potential role for $G_{\delta_{\nu}}$ in receptor-catalysed nucleotide exchange. In addition, the presence of receptor contact sites on both G_α and $G_{\beta\gamma}$ makes the stability of the receptor-G protein complex dependence dependent on the interaction between the subunits, and hence provides a simple mechanism for coupling the GTP-triggered dissociation of G_{α} and G_{β} , to their release from the receptor.

Ultimately, a mechanistic understanding of the nucleotideexchange process will require the structure of an activated heptahelical receptor bound to the nucleotide free form of a heterotrimeric G protein. In the interim, the present study provides the basis for mutational experiments designed to test specific hypotheses regarding the mechanism of nucleotide exchange and better define the role of $G_{\beta\gamma}$ in facilitating this process.

Note added in proof: After this paper was submitted, Wall et al. (Cell 83, 1047–1058, 1995) published a partially refined structure of $G_{ix1\beta1y2}$ showing a similar overall architecture. However, a comparison of the intersubunit contacts of the G, heterotrimer with those of the G_i heterotrimer is compromised by an admitted lack of 'atomicity' in the $G_{i\alpha 1\beta 1\gamma 2}$ model.

Received 1 December; accepted 18 December 1995

- 1. Conklin, B. R. & Bourne, H. R. Cell 73, 631-641 (1993).
- Neer, E. J. Cell 80, 249-257 (1995).
- 3. Simon, M. I., Strathmann, M. P. & Gautam, N. Science 282, 802-808 (1991).
- Hepler, J. R. & Gilman, A. G. Trends biochem. Sci. 17, 383-387 (1992).
 Crespo, P., Xu, N., Simonds, W. F. & Gutkind, J. S. Nature 369, 418-420 (1994).
- 6. Faure, M., Voyno-Yasenetskaya, A. & Boume, H. R. J. biol. Chem. 269, 7851-7854 (1994).
- Hurley, J. B. J. Bioenerg. Biomem. 24, 219–226 (1992).
- 8. Pfister, C. et al. Cell Sig. 5, 235-241 (1993).
- 9. Lambridght, D. G., Noel, J. P., Hamm, H. E. & Sigler, P. B. Nature 369, 621-628 (1994).
- Noel, J. P., Hamm, H. E. & Sigler, P. B. Nature 366, 654-663 (1993).
 Sondek, J. S., Lambright, D. G., Noel, J. P., Hamm, H. E. & Sigler, P. B. Nature 372, 276-279
- 12. Coleman, D. E. et al. Science **265**, 1405–1412 (1994). 13. Skiba, N. et al. J. biol. Chem. **271**, 413–424 (1996).
- 14. Mixon, M. B. et al. Science 270, 954 (1995). (initial?)
- 15. Neer, E. J., Schmidt, C. J., Nambudripad, R. & Smith, T. F. Nature **371**, 297–300 (1994). 16. Fung, B. K. & Nash, C.R. J. biol. Chem. **258**, 10503–10510 (1983).
- Mazzoni, M.R. & Hamm, H. E. Biochemistry 28, 9873–9880 (1989).
- 18. Mazzoni, M. R., Malinski, J. A. & Hamm, H. E. J. biol. Chem. 266, 14072-14081 (1991)
- 19. Graf, R., Mattera, R., Codina, J., Estes, M. K. & Birnbaumer, L. J. biol. Chem. 267, 24307-24314 (1992).
- 20. Denker, B. M., Neer, E. J. & Schmidt, C. J. biol. Chem. 267, 6272-6277 (1992).
- Slepak, V. Z., Wilkie, T. M. & Simon, M. I. J. biol. Chem 288, 1414–1423 (1993).
 Miller, R.T. et al. Nature 267, 712–715 (1988).
- Lee, E., Taussig, R. & Gilman, A. G. J. biol. Chem. 267, 1212–1218 (1992).
 Garcia-Higuera, I., Thomas, T. C., Yi, F. & Neer, E. J. J. biol. Chem. 271, 528 (1996)
- 25. Whiteway, M., Clark, K. L., Leberer, E., Dignard, D. & Thomas, D. Y. Molec. cell. Biol. 14, 3223-3229 (1994)
- 26. Sondek, J., Bohm, A., Lambright, D. G., Hamm, H. E. & Sigler, P. B. Nature 379, 369-374
- 27. Higashijima, T., Ferguson, K. M., Sternweis, P. C., Murray, D. S. & Gilman, A. G. J. blol. Chem. 262, 762-766 (1987).
- 28. Fukada, Y. et al. Nature 346, 658-660 (1990).
- 29. Lai, R. K., Dolores, P.-S., Canada, F. J. & Rando, R. R. Proc. natn. Acad. Sci. U.S.A. 87, 7673-
- 30. Mumby, S. M., Casey, P. J., Gilman, A. G., Gutowski, S. & Stemweis, P. C. Proc. natn. Acad. Sci. U.S.A. 87, 5873-5877 (1990).

- 31. Yamane, H. K. et al. Proc. natn. Acad. Sci. U.S.A, 87, 5868-5872 (1990).
- Maltese, W. A. & Robishaw, J. D. J. biol. Chem. 265, 18071–18074 (1990).
 Buss, J. E., Mumbry, S. M., Casey, P. J., Gilman, A. G. & Sefton, B. M. Proc. natn. Acad. Sci. U.S.A. 84, 7493-7497 (1987).
- 34. Kokame, K., Fukada, Y., Yoshizawa, T., Takao, T. & Shimonishi, Y. Nature 359, 749-752 (1992).
- 35. Neubert, T. A., Johnson, R. S., Hurley, J. B. & Walsh, K. A. J. biol. Chem. 267, 18274-18277 (1992)
- 36. Linder, M. E. et al. Proc. natn. Acad. Sci. U.S.A. 90, 3675-3679 (1993)
- 37. Wedegaertner, P. B. & Bourne, H. R. Cell 77, 1063-1070 (1994).
- 38. Linder, M. E. et al. J. blol. Chem. 266, 4654-4659 (1991)
- 39. Iniguez-Lluhi, J. A., Simon, M. I., Robishaw, J. D. & Cilman, A. G. J. biol. Chem. 267, 23409-
- 40. Bigay, J., Faurobert, E., Franco, M. & Chabre, M. Biochemistry 33, 14081-14090 (1994).
- 41. West, R. E., Moss, J., Vaughan, M., Liu, T. & Liu, T. Y. J. biol. Chem. 280, 14428—14430.
- 42. Sullivan, K. A. et al. Nature 330, 758-760 (1987).
- 43. Hirsch, J. P., Dietzel, C. & Kurjan, J. Genes Dev. 5, 467—474 (1991). 44. Simonds, W. F., Goldsmith, P. K., Codina, J., Unson, C. G. & Spiegel, A.M. Proc. natn. Acad. Sci.
- U.S.A. 86, 7809-7813 (1989).
- Conklin, B. R., Farfel, Z., Lustig, K. D., Julius, D. & Bourne, H. R. Nature 363, 274–276 (1993).
 Hamm, H. F. et al. Science 241, 832–835 (1988).
- 47. Dratz, E. A. et al. Nature **363**, 276-281 (1993). 48. Phillips, W. J. & Cerione, R. A. *J. biol. Chem.* **267**, 17032-17039 (1992)
- Stryer, L., Hurley, J. B. & Fung, B. K.-K. Meth. Enzym. 96, 617–627 (1983).
 Kabsch, W. J. appl. Crystallogr. 21, 67–71 (1988).
- Brunger, A. T. Acta crystallogr. A46, 46–57 (1990).
 Brunger, A. T. X-PLOR Version 3.1 Manual (Yale University, 1993).
- 53. Jones, T. A. et al. Acta crystallogr. A47, 110-1.19 (1991). 54. Grishin, A. V., Weiner, J. L. & Blumer, K. Molec. cell. Biol. **14**, 4571–4578 (1994). 55. Leberer, E. et al. EMBO J. **11**, 4805–4813 (1992).
- 56. Nicholls, A., Sharp, K.A. & Honig, B. Proteins 11, 281-296 (1991).

ACKNOWLEDGEMENTS, We thank M. Capel and L. Berman for access to and help with the X-25 beam line at the NSLS, J. Geiger for assistance during data collection, and E. Neer for making a manuscript available before publication. This work was supported by grants from the NIH to P.B.S. and H.E.H. H.E.H. was also supported by the American Heart Association, D.G.L. and J.S. were Damon-Runyon postdoctoral fellows. The coordinates of the heterotrimer structure will be submitted to the Protein Data Bank